Appl. No. 09/553,969 Amdt. dated September 30, 2011 Amendment under 37 CFR 1.116 Expedited Procedure Examining Group 1611

Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

1. (Previously Presented) A composition comprising:

an extrudable fragmented biocompatible resorbable single phase aqueous colloid which is substantially free from a free aqueous phase, said single phase aqueous colloid being present in an applicator having an extrusion orifice, wherein the single phase aqueous colloid has been fragmented by mechanical disruption, comprises a cross-linked gelatin polymer present in discrete subunits, has an equilibrium swell from 400% to 5000%, and has at least one characteristic selected from the group consisting of (a) a subunit size when fully hydrated in the range from 0.01 mm to 5 mm, and (b) an *in vivo* degradation time of less than one year;

wherein the single phase aqueous colloid is at least partially hydrated with an aqueous medium; and

wherein the aqueous medium comprises an active clotting agent that is thrombin.

- 2-18. (Canceled)
- 19. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid has a subunit size when fully hydrated in the range from 0.01 mm to 5 mm.
 - 20. (Canceled)
- 21. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid has an *in vivo* degradation time of less than one year.
 - 22-23. (Canceled)

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24. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid has a subunit size when fully hydrated in the range from 0.01 mm to 5 mm and an *in vivo* degradation time of less than one year.

25-29. (Canceled)

- 30. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid further comprises a polysaccharide.
- 31. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid further comprises a non-biological polymer.
- 32. (Previously Presented) The composition of claim 1, wherein the single phase aqueous colloid further comprises a polysaccharide or a non-biological polymer, or both.

33-34. (Canceled)

35. (Previously Presented) A composition comprising:

an extrudable fragmented biocompatible resorbable single phase aqueous colloid which is substantially free from a free aqueous phase, wherein the single phase aqueous colloid has been fragmented by mechanical disruption, has an equilibrium swell from 400% to 5000%, and comprises a cross-linked protein present in discrete subunits and a polysaccharide, the single phase aqueous colloid having at least one characteristic selected from the group consisting of (a) a subunit size when fully hydrated in the range from 0.01 mm to 5 mm and (b) an *in vivo* degradation time of less than one year;

wherein the single phase aqueous colloid is at least partially hydrated with an aqueous medium;

wherein the aqueous medium comprises an active clotting agent that is thrombin; and

wherein the cross-linked protein is present in an applicator having an extrusion orifice.

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36. (Previously Presented) A composition comprising:

an extrudable fragmented biocompatible resorbable single phase aqueous colloid which is substantially free from a free aqueous phase, wherein the single phase aqueous colloid has been fragmented by mechanical disruption, has an equilibrium swell from 400% to 5000%, and comprises a cross-linked protein present in discrete subunits and a non-biological polymer, the single phase aqueous colloid having at least one characteristic selected from the group consisting of (a) a subunit size when fully hydrated in the range from 0.01 mm to 5 mm and (b) an *in vivo* degradation time of less than one year;

wherein the single phase aqueous colloid is at least partially hydrated with an aqueous medium;

wherein the aqueous medium comprises an active clotting agent that is thrombin; and

wherein the cross-linked protein is present in an applicator having an extrusion orifice.

37. (Withdrawn) A device consisting of:

a syringe; and

an amount of a resorbable fragmented cross-linked gelatin gel present in the syringe, wherein the gel biodegrades in a patient's body in a time period ranging from 1 to 90 days.

- 38. (Withdrawn) The device according to Claim 37, wherein the gel biodegrades in a patient's body in a time period ranging from 2 to 30 days.
- 39. (Withdrawn) The device according to Claim 37, wherein the gel resorbs in a time period ranging from 14 to 60 days.
 - 40. (Withdrawn) A device consisting of:

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a syringe;

an amount of a resorbable fragmented cross-linked gelatin gel present in the syringe, wherein the gel biodegrades in a patient's body in a time period ranging from 1 to 90 days; and

a bioactive component.

- 41. (Withdrawn) The device according to Claim 40, wherein the bioactive component is a hemostatic agent.
- 42. (Withdrawn) The device according to Claim 41, wherein the hemostatic agent is thrombin.
- 43. (Withdrawn) The device according to Claim 42, wherein the gel comprises 500 to 1000 units thrombin/ml gel.
 - 44. (Withdrawn) A composition of matter comprising:
 - a sterile package; and
 - a device according to Claim 37 present inside of the sterile package.
 - 45. (Withdrawn) A device consisting of:
 - a syringe; and

an amount of a resorbable fragmented partially hydrated cross-linked gelatin gel present in the syringe.

46. (Withdrawn) The device according to Claim 45, wherein the gel has an equilibrium swell ranging from 400% to 1300%.

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- 47. (Withdrawn) The device according to Claim 46, wherein the gel has an equilibrium swell ranging from 500% to 1100%.
- 48. (Withdrawn) The device according to Claim 47, wherein the gel biodegrades in a patient's body in a time period ranging from 1 to 90 days.
- 49. (Withdrawn) The device according to Claim 45, wherein the gel biodegrades in a patient's body in time period ranging from 2 to 30 days.
- 50. (Withdrawn) The device according to Claim 45, wherein the gel resorbs in a time period ranging from 14 to 60 days.
 - 51. (Withdrawn) A device consisting of: a syringe; and

an amount of a resorbable fragmented cross-linked gelatin gel present in the syringe wherein the gel has an equilibrium swell from 400% to 1300%.

- 52. (Withdrawn) The device according to Claim 51, wherein the gel has an equilibrium swell ranging from 500% to 1100%.
- 53. (Withdrawn) The device according to Claim 51, wherein the gel biodegrades in a patient's body in a time period ranging from 1 to 90 days.
- 54. (Withdrawn) The device according to Claim 51, wherein the gel biodegrades in a patient's body in time period ranging from 2 to 30 days.
- 55. (Withdrawn) The device according to Claim 51, wherein the gel resorbs in a time period ranging from 14 to 60 days.

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- 56. (Withdrawn) The device according to Claim 51, wherein the gel comprises a bioactive component.
- 57. (Withdrawn) The device according to Claim 56, wherein the bioactive component is a hemostatic agent.
- 58. (Withdrawn) The device according to Claim 56, wherein the hemostatic agent is thrombin.
- 59. (Withdrawn) The device according to Claim 58, wherein the gel comprises 100 to 1000 units thrombin/ml gel.
 - 60. (Withdrawn) A kit comprising: a device according to either Claim 45 or Claim 51; and a tray.
- 61. (Withdrawn) The kit according to Claim 60, wherein the kit further comprises a container comprising an aqueous medium.
- 62. (Withdrawn) The kit according to Claim 61, wherein the kit further comprises thrombin.
 - 63. (Withdrawn) A method comprising:
 - (a) providing a device consisting of:
 - (i) a syringe; and

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(ii) an amount of a resorbable fragmented cross-linked gelatin gel present in the syringe, wherein the gel biodegrades in a patient's body in a time period ranging from 1 to 90 days; and

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(b) delivering the gel from the syringe to a patient.